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Abstract: Non-invasive detection of specific atherosclerotic plaque components related to vulnerability is of high clinical relevance to prevent cerebrovascular events. The feasibility of magnetic resonance imaging (MRI) for characterization of plaque components was already demonstrated. We aimed to evaluate the potential of ex vivo differential phase contrast X-ray tomography (DPC) to accurately characterize human carotid plaque components in comparison to high field multicontrast MRI and histopathology. Two human plaque segments, obtained from carotid endarterectomy, classified according to criteria of the American Heart Association as stable and unstable plaque, were examined by ex vivo DPC tomography and multicontrast MRI (T1-, T2-, and proton density-weighted imaging, magnetization transfer contrast, diffusion-weighted imaging). To identify specific plaque components, the plaques were subsequently sectioned and stained for fibrous and cellular components, smooth muscle cells, hemosiderin, and fibrin. Histological data were then matched with DPC and MR images to define signal criteria for atherosclerotic plaque components. Characteristic structures, such as the lipid and necrotic core covered by a fibrous cap, calcification and hemosiderin deposits were delineated by histology and found with excellent sensitivity, resolution and accuracy in both imaging modalities. DPC tomography was superior to MRI regarding resolution and soft tissue contrast. Ex vivo DPC tomography allowed accurate identification of structures and components of atherosclerotic plaques at different lesion stages, in good correlation with histopathological findings.

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Ex vivo Differential Phase Contrast and Magnetic Resonance Imaging for Characterization of Human Carotid Atherosclerotic Plaques

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Abstract

Aims

Non-invasive detection of specific atherosclerotic plaque components related to vulnerability is of high clinical relevance to prevent cerebrovascular events. The feasibility of magnetic resonance imaging (MRI) for characterization of plaque components was already demonstrated. We aimed to evaluate the potential of ex vivo differential phase contrast X-ray tomography (DPC) to accurately characterize human carotid plaque components in comparison to high field multicontrast MRI and histopathology.

Methods and Results

Two human plaque segments, obtained from carotid endarterectomy, classified according to criteria of the American Heart Association as stable and unstable plaque, were examined by ex vivo DPC tomography and multicontrast MRI (T1-, T2-, and proton density-weighted imaging, magnetization transfer contrast, diffusion-weighted imaging). To identify specific plaque components, the plaques were subsequently sectioned and stained for fibrous and cellular components, smooth muscle cells, hemosiderin, and fibrin. Histological data were then matched with DPC and MR images to define signal criteria for atherosclerotic plaque components. Characteristic structures, such as the lipid and necrotic core covered by a fibrous cap, calcification and hemosiderin deposits were delineated by histology and found with excellent sensitivity, resolution and accuracy in both imaging modalities. DPC tomography was superior to MRI regarding resolution and soft tissue contrast.

Conclusion

Ex vivo DPC tomography allowed accurate identification of structures and components of atherosclerotic plaques at different lesion stages, in good correlation with histopathological findings.

Key Words

X-rays, Synchrotron, Differential phase contrast, Magnetic resonance imaging, Carotid plaque, Atherosclerosis

Abbreviations

DPC=differential phase contrast, MRI=magnetic resonance imaging, PD=proton density, MTC=magnetization transfer contrast, DW=diffusion-weighted, L=lumen of the blood vessel, M=media, Int=intima, IT=intimal thickening, FC=fibrous cap, LC=lipid core, NC=necrotic core, U=ulceration site.

Introduction

The rupture of atherosclerotic plaques in carotid arteries is the main cause for stroke. Every year, 15 million people suffer a stroke worldwide. One-third of the affected die, one-third are left permanently disabled and one-third will recover completely from their attack¹. Projections show that by the year 2030, an additional 3.4 million people aged ≥ 18 years will have had a stroke corresponding to 20.5% increase in prevalence from 2012². To prevent stroke, the most frequently performed surgical intervention is carotid endarterectomy.

Most ruptures occur in so called unstable plaques containing a highly inflamed lipid-rich and/or necrotic core covered by a thin fibrous cap³. Early identification and characterization of plaques are crucial for risk prediction and prevention of adverse events. Invasive techniques, such as intravascular ultrasound, optical coherent tomography, and carotid angiography can reveal luminal stenosis, wall thickness and plaque volume. However, plaque rupture and stroke are not dependent on the extent of stenosis. Characterizing plaque morphology and, in particular, plaque stabilizing and destabilizing components may provide more information on plaque instability and consequently on the risk of rupture. Besides molecular imaging such as positron emission tomography (PET) and single photon emission computed tomography (SPECT), the two imaging techniques computed tomography (CT) and magnetic resonance imaging (MRI) that are widely accepted by the medical community, have the potential to assess plaque vulnerability non-invasively. However, to detect density differentials within soft tissues, such as in atherosclerotic plaques, a good sensitivity coupled with high-resolution are required, which does not apply to conventional CT and MRI. A recently introduced technique, synchrotron-based differential phase contrast X-ray (DPC) tomography⁴⁻⁶, does not only measure the attenuation of X-rays passing through tissue, as in standard X-ray tomography, but in addition it measures the phase shift caused by refraction, which the coherent X-ray beam undergoes when passing through tissue with different refractive indices. This results in a high sensitivity to electron density differences and thereby improving contrast for soft tissue. The objective of our study was to evaluate the feasibility of ex vivo DPC tomography to morphologically

characterize two human carotid atherosclerotic plaque specimens. Multicontrast-weighted MRI is capable of imaging vessel wall structures and carotid plaque compositions *in vivo*⁷ and *ex vivo*⁸. MRI differentiates plaque components on the basis of water content, physical state, and molecular motion or diffusion while providing information-rich images at high spatial resolution. The most common strategy is to analyze multicontrast images (T1-, T2-, proton density (PD)-weighted imaging) that reveal components with different contrast⁹. The content and distribution of lipids is an important aspect of atherosclerotic plaques. To improve the contrast between lipids and fibrous tissue, the low mobility of protons bound to lipids has been exploited by using diffusion-weighted (DW) imaging¹⁰. For better identification of protein-rich regions magnetization transfer contrast (MTC) was recommended by Qiao et al.¹¹.

To our knowledge no direct comparison of DPC images, multicontrast MR images and histopathology of human endarterectomized carotid plaques has been carried out. The purpose of this *ex vivo* study was (i) to explore the potential of a grating interferometer at a synchrotron X-ray source to measure the DPC between different components in human stable and unstable carotid plaques, (ii) to compare the images with multicontrast high-resolution MRI and (iii) to correlate the generated images from both modalities with histopathology.

Methods

Study design and experimental overview

Written informed consent was obtained from a 75-year-old male patient scheduled for carotid endarterectomy. Two carotid specimens, one originating from the *Arteria carotis externa* and the other from the *Arteria carotis communis/interna*, were immediately transferred into RNAlater® (Sigma, St. Louis, USA) solution and stored at 4°C overnight according to our standardized procedure for plaque harvesting and bio-banking¹². The next day, tissues were fixed in 4% formalin for 24 h at 4°C and finally stored in PBS at 4°C.

First, the two specimens were scanned by DPC tomography at the TOMCAT (Tomographic Microscopy and Coherent Radiology Experiments) beamline of the Swiss Light Source, followed by imaging on a small animal MR system using different MR contrasts. In a second step, both plaques were characterized by histological workup and, finally, the DPC and MR images were matched with the corresponding histological sections.

Differential phase contrast imaging and three-dimensional rendering

The presented data sets were obtained by synchrotron-based X-ray tomography using the differential phase contrast technique^{4,6} at a photon energy of 25 keV. Carotid artery segments were placed in a falcon tube filled with PBS. To measure the phase shift, a grating interferometer was used and series of projection images were taken while the carotid plaque specimens rotated. A 300 μm thick LAG:Ce scintillator converted the X-rays to visible light, which were captured by a CMOS camera (PCO.Edge, PCO AG, Kelheim, Germany). The total scan time for a segment with a field of view (FOV) of 12 mm x 3.5 mm was about 90 min. More details about components and parameters are provided in Table S1.

To obtain 3-dimensional images of the DPC data sets, individual Tiff files were converted to a multilayer Tiff file and processed by Imaris software (Bitplane AG, Zurich, Switzerland).

Magnetic resonance imaging

MRI was performed with a Bruker Pharmascan 7/16 small animal MR system equipped with a gradient system capable of a maximum gradient strength of 760 mT/m, with a 80 μs rise time and a quadrature birdcage resonator. The specimens were placed in a Falcon tube, filled with PBS, inside the volume resonator and kept at 24°C. Reference data was acquired in coronal and sagittal orientations for accurate positioning of the plaque specimen. Before imaging a fieldmap-based local shimming was performed on the specimen using the automated MAPshim routine to reduce field inhomogeneities. The imaging protocol consisted of different spin echo (SE) sequences using

different dimensions of the FOVs, matrix sizes, echo times (TE) and repetition times (TR) as shown in Table S2. For all sequences 30 averages were performed.

To calculate the MTC, images with (MT_{on}) and without (MT_{off}) the application of a saturation pulse were acquired¹¹. For MT offset the frequency and amplitude were optimized and a Gaussian saturation pulse with an offset frequency of 3.5 kHz and amplitude of 30 μ T was found optimal. MT subtraction maps were calculated using the equation:

$$MTC = MT_{on} - MT_{off}.$$

For DW imaging, diffusion-encoding was applied (gradient pulse duration = 2.5 ms, gradient pulse separation = 8.1 ms) with a b-value of 650 s/mm^2 .

Histological processing and histopathology

After DPC and MRI examinations, the two carotid plaques were paraffin-embedded and serial sections of 2.5 μ m were prepared for further histological and immunohistochemical investigations. Sections were routinely stained with hematoxylin and eosin (HE), Masson's trichrome, Elastica van Gieson (VG-Elastica), phosphotungstic acid hematoxylin (PTAH) showing fibrin deposits in lesions, and Prussian blue staining to identify iron-containing hemosiderin from previous hemorrhage. For immunohistochemistry, the monoclonal antibody anti-human alpha smooth muscle cell actin (anti-SMA, 1:400, mouse, M0851, Dako, Baar, Switzerland) was used. The detection system included the Dako RealKit (Dako) on the immunostainer (Dako). All sections were digitized by a slide scanner with a pixel size of 0.221 x 0.221 μ m² (Pannoramic 250, 3D Histech, Sysmex, Horgen, Switzerland). Pathological classification of the two plaque types was done in a first instance macroscopically by the surgeon (Z.R.) according to their surface morphology and further investigated on the basis of the modified American Heart Association (AHA)-classifications¹³ by a board-certified pathologist (N.B.) using histology and specific staining methods. The experienced pathologist (N.B.) identified vessel layers and plaque components, such as media (M), intima (Int), intimal thickening (IT), fibrous cap (FC), lipid core (LC), necrotic core (NC), inflammatory cell infiltration, hemosiderin deposits, neovascularization, and calcifications.

Interpretation of DPC and MRI data

The DPC images, MR images and histological slides were matched using the known location and distance between DPC images, MR images and histological cross sections. Furthermore, gross morphological features, such as plaque shape, vessel wall thickness and shape as well as calcium deposits were used to optimize the matches. We did not account for shrinkage of the specimen caused by histological processing, as it can vary across the specimen and would require multiple landmarks for accurate matching¹⁴. The above mentioned plaque components were identified in both imaging modalities using the histological sections as reference. Three independent investigators (R.M., P.S., A.M.H.) analyzed DPC and MR images for plaque signal characteristics in comparison with surrounding structures.

Results

Morphological characterization of a stable and unstable plaque by histology

According to histopathology, the specimen originating from the *Arteria carotis externa* was classified as a stable plaque (Figure 1A) and the specimen obtained from *Arteria carotis communis/interna* as an unstable plaque (Figure 1B).

Histological workup was performed at the end of the ex vivo imaging procedures. Figure 2 shows the histological identification of the stable and unstable lesions. With HE, the investigated stable plaque presented a thick fibrous cap with fibrillar eosinophilic material. A high collagen content (blue) and local accumulation of erythrocytes (red) between the fibrous cap and the intima was proven by Masson's trichrome staining. A staining for iron-containing hemosiderin (Prussian blue) was negative. Elastic fibers, stained black with VG-Elastica, were present in the lamina elastica interna and media. Immunohistochemical analyses using an anti-SMA antibody demonstrated the presence of smooth muscle cells (SMC) in the lamina elastica interna and media.

The unstable plaque presented towards one plaque shoulder a large necrotic core overlaid by a thicker fibrous cap and towards the other plaque shoulder a smaller lipid core overlaid by a thinner fibrous cap. Invasion of many macrophages and immune cells into the lipid core and partially necrotic core was strongly indicated by HE. The necrotic core contained cell debris, necrotic material and cholesterol crystals. A locally extensive calcification (arrow) was clearly detected with HE. Erythrocytes in the lipid core were of intensive red and SMCs in the media were stained light red by Masson's trichrome. Intracellular hemosiderin pigments, a sign of erythrophagocytosis indicating preceding plaque hemorrhage, were found in macrophages in the lipid core (Figure 2a high-power magnification, Prussian blue). Positive Prussian blue staining in the media (Figure 2b, high-power magnification) is linked to the ability of SMCs to phagocytose heme-derived iron^{15, 16}. The staining of SMCs correlated with the presence of elastic fibers in the media and lamina elastica interna. Few SMCs were also present in regions of the fibrous cap. PTAH staining identified intraplaque fibrin in the lipid core and necrotic core of the unstable plaque. Fibrin deposition in lipid core/necrotic core regions are known to delineate late stage plaques¹⁷. Furthermore PTAH staining revealed a ruptured fibrous cap as visualized by a fibrin-positive ulceration site (U) towards the lumen of the blood vessel.

Appearances of atherosclerotic plaque components in DPC tomography and multicontrast MRI

In the following section we demonstrate the similarities and differences in appearance of plaque components among the different imaging modalities and in correlation with histopathology. Figure 3A and Figures S1 and S2 show the matched DPC images, MR images and histopathology sections of the stable and unstable carotid plaque. Visual appearances of each plaque component in DPC and MR tomography are summarized in Table 1. The high-resolution and sensitivity of DPC tomography enabled high-power magnification images which are shown with the matching HE stained sections in Figure 3B-K. Animated GIF-files of the DPC scans are provided in Video S1 and S2.

Tunica media

Histopathology revealed SMCs and elastic fibers in the media of the stable and unstable plaque. The stable plaque displayed a media of homogeneous texture in DPC and MR images (Figure 3A, C). Hyperintense (bright signal) T1-weighted and DW images distinguished the media from the intima. In contrast, media of the unstable plaque displayed hypointense (dark signal) T1-, T2-, DW-, and PD-weighted images, hyperintense MTC images, and an isodense signal in DPC images. Staining for hemosiderin disclosed blue precipitates as iron-containing particles engulfed by SMCs in the media of the unstable plaque which account for prominent signal intensities.

Tunica intima

The stable plaque consisted of an intact intima, with a homogeneous texture and intermediate intensity in most image contrasts, except for MTC and DW images where they appeared hypointense and hyperintense, respectively (Figure 3A, C).

Intimal thickening

The intimal thickening in the stable plaque appeared homogenous and hyperdense in DPC delimiting it from the intima. MTC images revealed a hypointense intimal thickening (Figure 3A, D).

Erythrocyte accumulation

The gap between intima and intimal thickening/fibrous cap of the stable plaque was clearly defined in DPC and T1 contrast, but less in the other MR settings (Figure 3A). The region was appreciated as low homogenous signal with a cloudy appearance in the zoomed DPC image (Figure 3E). The presence of erythrocytes was only proven by histological staining.

Fibrous cap

The fibrous cap of both the stable and unstable plaque showed homogeneous and intermediate intensities in DPC images and MR images, except in MTC where it appeared hypointense (Figure 3A, F).

Lipid core

A lipid core was only visible in the unstable plaque composed of hemosiderin-containing macrophages and erythrocytes as well as fibrin. The lipid core appeared hypointense in T1-, T2-, DW-, and PD-weighted images, and hyperintense in MTC (Figure 3A). The lipid core could be delineated

from intima and fibrous cap in all setups. The lipid core had an inhomogeneous texture from isodense to hypodense in DPC (Figure 3G).

Necrotic core

The necrotic core was clearly visible in all MR images with varying signal intensities, however, with more homogenous appearance than in DPC (Figure 3A). In hypodense areas of the DPC images we identified cholesterol crystals or dissolved crystals and in hyperdense regions cell debris were identified (Figure 3H and I, respectively).

Calcifications

The highest and distinct signal intensity of all plaque components was found in regions of calcification in the unstable plaque (Figure 3A, J). High-dense calcification (high amount of calcium, arrow) appeared hyperdense/-intense in DPC and MTC. In T1-, T2-, DW- and PD-weighted images it appeared hypointense. This calcification was sharply delineated from the surrounding soft tissue. In contrast, low-dense calcification (arrowhead) was distinguished only on T1-weighted images as hyperintense.

Ulceration

The focal ulceration site in the unstable plaque was fibrin-positive and appreciated as a ruptured cap in HE and DPC (Figure 3K) and was not clearly discernable in MR images.

Three-dimensional visualization of endarterectomized atherosclerotic plaques in DPC

The three-dimensional rendering of DPC images of the carotid plaque specimens are shown in Figure 4. The needle-shaped crystal in the unstable plaque appeared, in the given field of view, with an approximate length of 1.5 mm (Figure 4E). Due to the limited field of view in DPC tomography only a part of the calcified crystal was imaged, however, allowing estimating that a relatively large calcification was present in this plaque specimen.

Discussion

In this ex vivo feasibility study we were able to demonstrate that detailed information on morphological characteristics and composition of a human stable and unstable endarterectomized plaque can be obtained with DPC tomography and multicontrast MRI. Our observations are in good correlation with histopathology.

Assessment of anatomy and composition of atherosclerotic plaques by MRI have been extensively studied ex vivo^{8, 18-21} and in vivo^{7, 22-24} using animal models of disease and in patients. However, a comprehensive evaluation of different MRI sequences has not been determined. With our combination of high-resolution MRI sequences, we were able to accurately identify fibrocellular tissue, lipid-rich and calcified regions as well as iron-containing deposits in the atherosclerotic plaques. Our intention was to apply the findings of MR images and histopathology to understand appearances of plaque characteristics in DPC images. In general, the DPC images presented a lower contrast than MR images, however, homo- and heterogeneous textures within the plaques are more pronounced in DPC images. All fibrocellular tissue, e.g. media, fibrous cap and intimal thickening, was recognized as a homogeneous pattern and delimited to the plaque. Regions of active remodeling within the unstable plaque (lipid/necrotic core) were clearly recognized as areas with heterogeneous appearance and different signal intensities. High-dense, but not low-dense, calcification gave a distinct hyperdense signal that clearly allowed for differentiation from background structures. Contrary to MRI, hemosiderin deposits presented no significant signal in DPC. There are a few studies²⁵⁻²⁸ reporting on the application of phase contrast imaging to atherosclerotic vessels and findings are in good agreement with our results.

With our study we provided a foundation for directing and validating the interpretation of MR and DPC images and attempted to develop and optimize imaging parameters. The relatively new technique, DPC tomography was conducted at a synchrotron radiation facility, which, so far, cannot be used in clinical practice and limits it to a benchmarking feasibility work. Recording one tomogram with the experimental parameters used in this study would result in a deposition of approximately

30'000 Gy⁴. For humans, a whole-body irradiation of up to 1 Gy is unlikely to cause long range symptoms, whereas a dose of > 30 Gy is always fatal²⁹. MRI has distinct advantages over DPC including no ionizing radiation and it can be applied in vivo³⁰ and hence to e.g. monitor plaque progression in longitudinal studies.

There are a number of limitations to this proof-of-concept study. First, the low sample number of only one stable and one unstable carotid plaque prevent us to raise quantitative data about the sensitivity, specificity and accuracy of DPC tomography and MRI. Second, assessment of sensitivities for different lesion components in DPC and MRI was done retrospectively. For further studies, the morphological appearances of lesion components in DPC or MRI, as listed in Table 1, need to be tested for suitability in the detection and differentiation of plaque characteristics in a prospective way. Third, the natural postmortem degradation processes of the endarterectomized plaques have to be taken into account including shrinkage during histological processes. Forth, in spite of technological progress, DPC tomography currently remains an experimental method. Further improvements are necessary to achieve an X-ray source with acceptable radiation doses and acquisition time before applying to humans.

In conclusion, our study demonstrates that DPC tomography can produce remarkable high-resolution images and can discriminate between clinically relevant components of the atherosclerotic vessel wall. Once adequately validated and optimized, DPC tomography might potentially help to define high-risk atherosclerotic plaques. Finally, we want to emphasize the importance of multicontrast imaging since only one contrast cannot differentiate all components. However, using multicontrast MRI will also not allow discriminating each single component, e.g. calcification and iron deposits displayed the same MR signal. More sophisticated diagnostic tools (e.g. new MR protocols) and validation studies will help understanding different contrasts.

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Figures

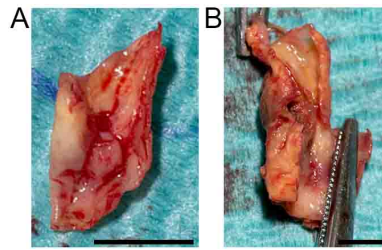


Figure 1 Whole mount human endarterectomized carotid plaques. (A) Stable plaque excised from *Arteria carotis externa*. (B) Unstable plaque excised from *Arteria carotis communis/interna*. Scale bars: 0.5 cm.

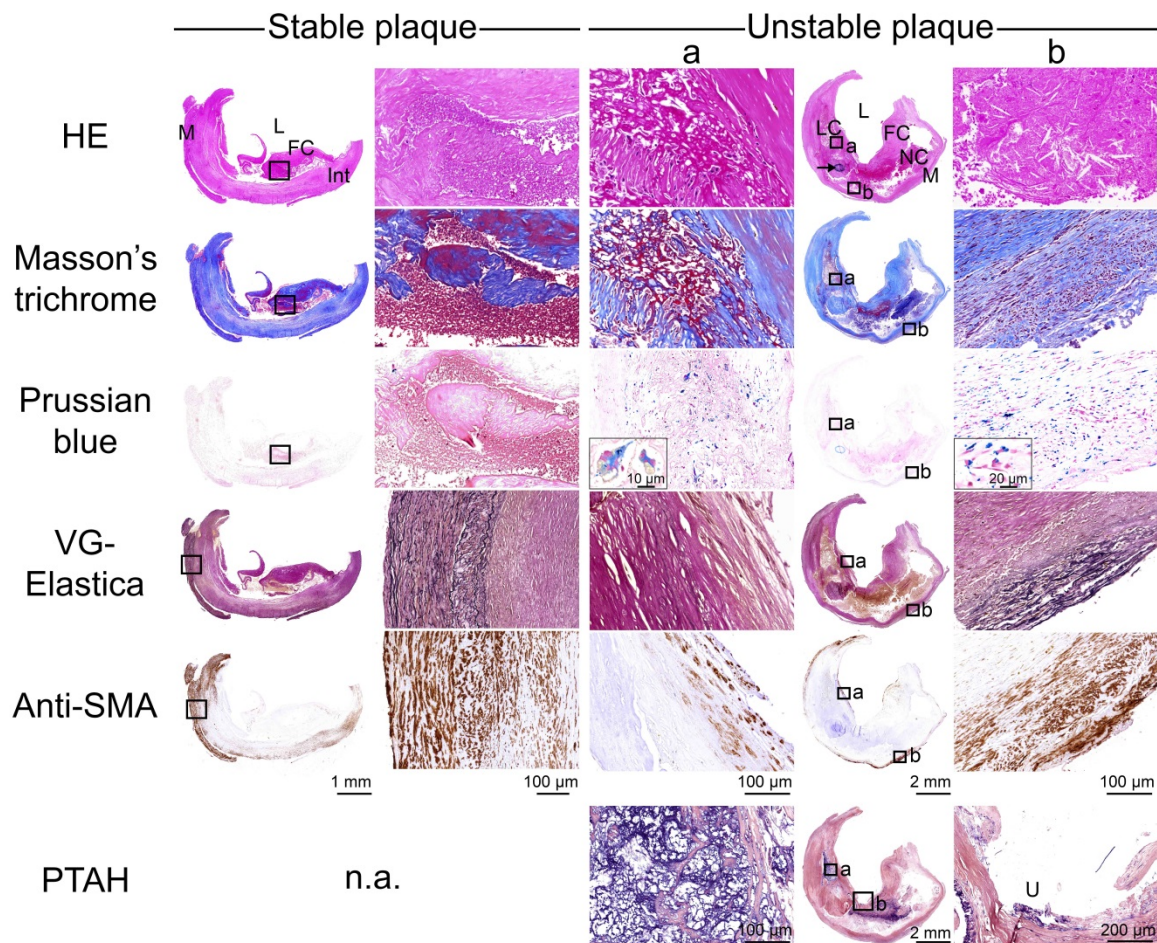


Figure 2 Histopathological sections of the stable and unstable plaque stained with hematoxylin and eosin (HE); Masson's trichrome coloring connective tissue in blue, erythrocytes and lipid/necrotic core in intense red, smooth muscle cells in light red; Prussian blue showing iron-containing hemosiderin (blue) indicating preceding intraplaque hemorrhage; Van Gieson-Elastica (VG-Elastica) coloring lamina elastica interna and other elastic fibers in black; anti-smooth muscle cell actin (anti-SMA) antibody labeling smooth muscle cells in media and fibrous cap. Phosphotungstic acid hematoxylin staining (PTAH) showing fibrin in purple located in lipid and necrotic core as well as at the ulceration site. Arrow in HE-stained unstable plaque section pointing at an extensive calcification. Boxed higher magnification images of the Prussian blue stained unstable plaque show hemosiderin-loaded macrophages present in the lipid core (a) and hemosiderin-loaded smooth muscle cells present in the media (b). L, lumen of the blood vessel; M, media; Int, intima; FC, fibrous cap; LC, lipid core; NC, necrotic core; U, ulceration site; n.a., not applicable.

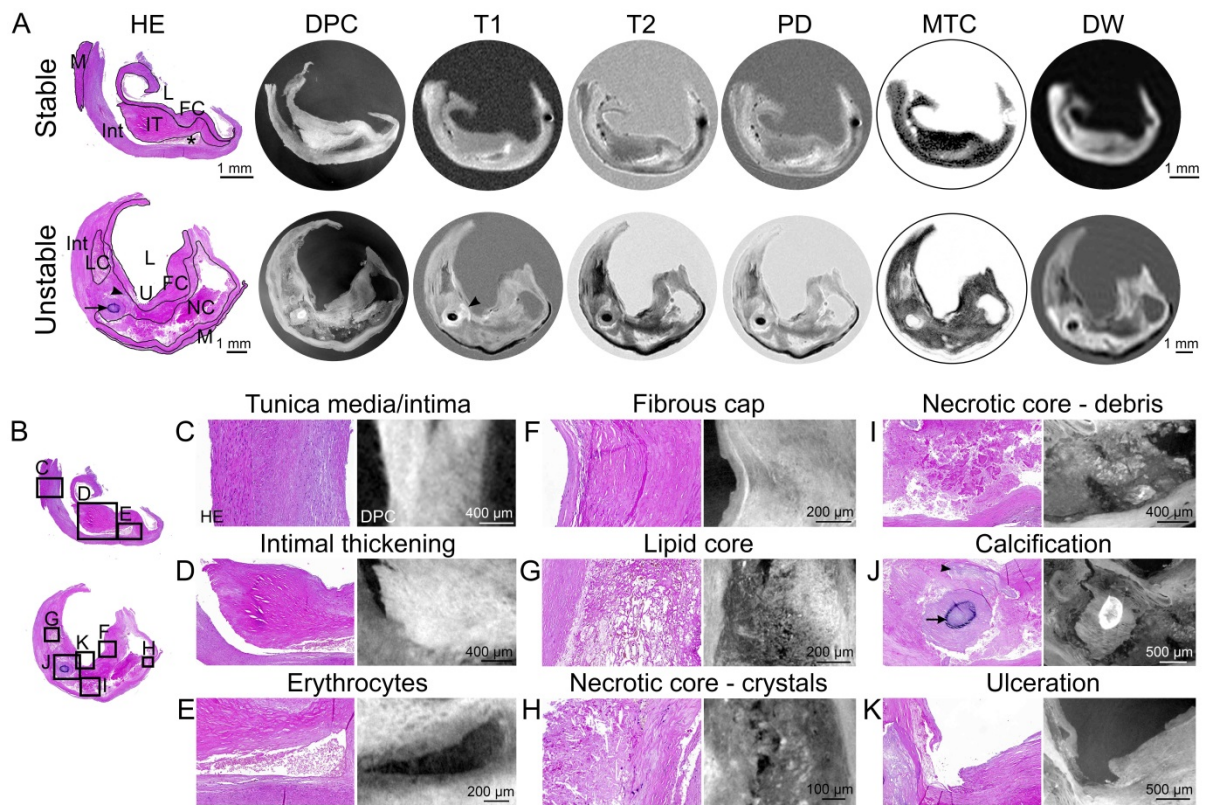
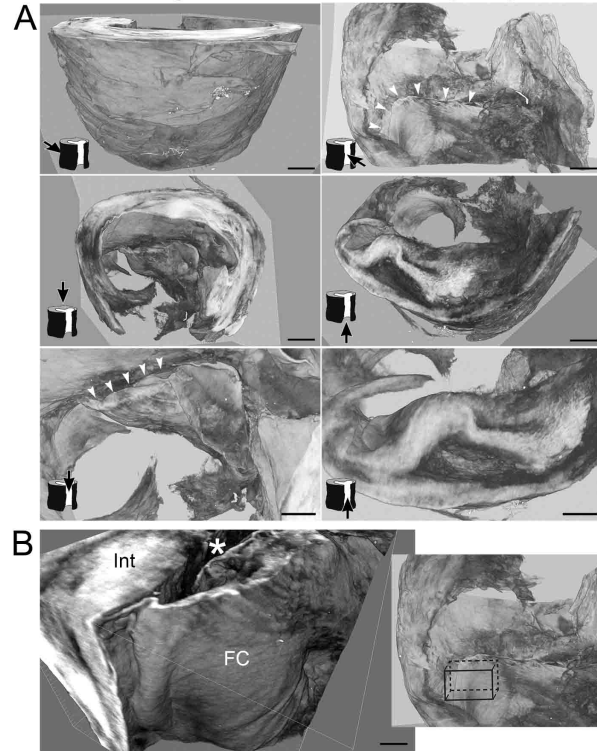


Figure 3 (A) Representative set of matched images of hematoxylin and eosin staining (HE), differential phase contrast (DPC) tomography and multicontrast MR (T1-, T2-, and proton density (PD)-weighted images, magnetization transfer contrast (MTC), diffusion-weighted (DW) images) of the stable and unstable plaque. For orientation, lumen of the blood vessel (L), media (M), intima (Int), intimal thickening (IT), fibrous cap (FC), lipid core (LC), necrotic core (NC), ulceration site (U) are labelled. Asterisk in stable plaque indicates erythrocyte accumulation; arrowhead in unstable plaque indicates low-dense calcifications; arrow indicates high-dense calcification. (B) Overview section of the stable (top) and unstable (bottom) plaque with boxed regions of interest for high-power magnification images in C-K. (C-K) Histopathology (HE-stained) and corresponding DPC tomograms of different plaque components of the stable (C-E) and unstable (F-K) plaque.

Carotid segment with stable plaque



Carotid segment with unstable plaque

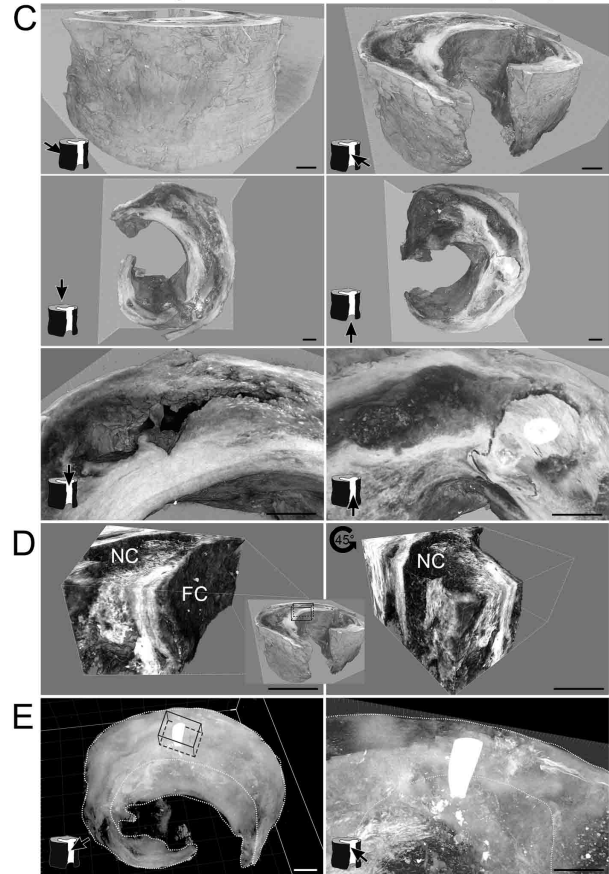


Figure 4 Three-dimensional DPC images of the endarterectomized plaques. Black schematic plaque with arrow indicates viewing direction. (A) Carotid segment with the stable plaque at different orientations, white arrowheads pointing to the plaque; scale bars: 3 mm. (B) Cropped region of the stable plaque showing intact and smooth surface of the fibrous cap (FC), intima (Int) and the cleft (asterisk) containing erythrocytes; scale bar: 1 mm. (C) Carotid segment with the unstable plaque at different orientations; scale bars: 1 mm. (D) Cropped region of the unstable plaque showing thin fibrous cap (FC) covering the necrotic core (NC); scale bars: 0.5 mm. (E) Left image showing the whole unstable plaque in a maximal intensity projection. Right image showing cropped region with calcified crystal; scale bars: 1 mm.

Tables

Table 1 Morphologic characteristics of plaque components compared with appearance in DPC and MRI

Components	DPC	T1	T2	PD	MTC	DW
Media (stable)	Isodense	Hyperintense	Isointense	Isointense	Isointense	Hyperintense
Media with hemosiderin (unstable)*	Isodense	Hypointense	Hypointense	Hypointense	Hyperintense	Hypointense
Intima	Isodense	Hyper- to Isointense	Isointense	Isointense	Hypointense	Hyperintense
Intimal thickening	Hyperdense	Isointense	Isointense	Isointense	Hypointense	Isointense
Fibrous cap	Isodense	Isointense	Isointense	Isointense	Hypointense	Isointense
Lipid core	Iso- to hypodense	Iso- to hypointense	Hypointense	Iso- to hypointense	Hyperintense	Hypointense
Necrotic core	Hyper- to hypodense	Isointense	Hyper- to hypointense	Iso- to hypointense	Isointense	Hyper- to hypointense
Calcifications						
Low-dense	Isodense	Hyperintense	Isointense	Isointense	Isointense	Isointense
High-dense	Hyperdense	Hypointense	Hypointense	Hypointense	Hyperintense	Hypointense

* Media of the unstable plaque contained iron-containing hemosiderin engulfed by SMCs

attributable for intense signals in MRI.

Supplementary Data

Table S1 Components and parameters of the grating interferometer-based DPC experiment

Component or Parameter	Specification or Value
X-ray source	Synchrotron
Ring current	400 mA
Beam energy	25 keV
Monostripe	W/Si
Field of View	12 mm x 3.5 mm
Detector	Complementary Metal Oxide Silicon (CMOS)
Sample Magnification	1.0
Scintillator	LAG:Ce 300 μm
Pixel	2048 x 2048
Pixel size	6.5 x 6.5 μm^2
Grating interferometer	7 phase steps
Scan settings	
Projections	1441 over 180°
Number of darks	32
Number of flats	100
Angular step	0.125°

Table S2 MRI sequence parameters

Sequence	TE [ms]	TR [ms]	Acquisition time	Geometry
T1	6.5	810	1h 17 min	For T1, T2, PD, MT: Slice thickness: 0.5 mm
T2	30	2000	3h 12min	No slice gap FOV: 15 x 15 mm
Proton density (PD)	13	2000	3h 12min	Matrix: 256 x 256 In-plane resolution: 59 x 59 μ m
Magnetization transfer (MT _{on} , MT _{off})	5	750	1h 36min (each)	
Diffusion-weighted (DW)	17.5	2500	4h	For DW: Slice thickness: 0.8 mm Slice gap: 0.25 mm FOV: 25 x 25 mm Matrix: 128 x 128 In-plane resolution: 195 x 195 μ m

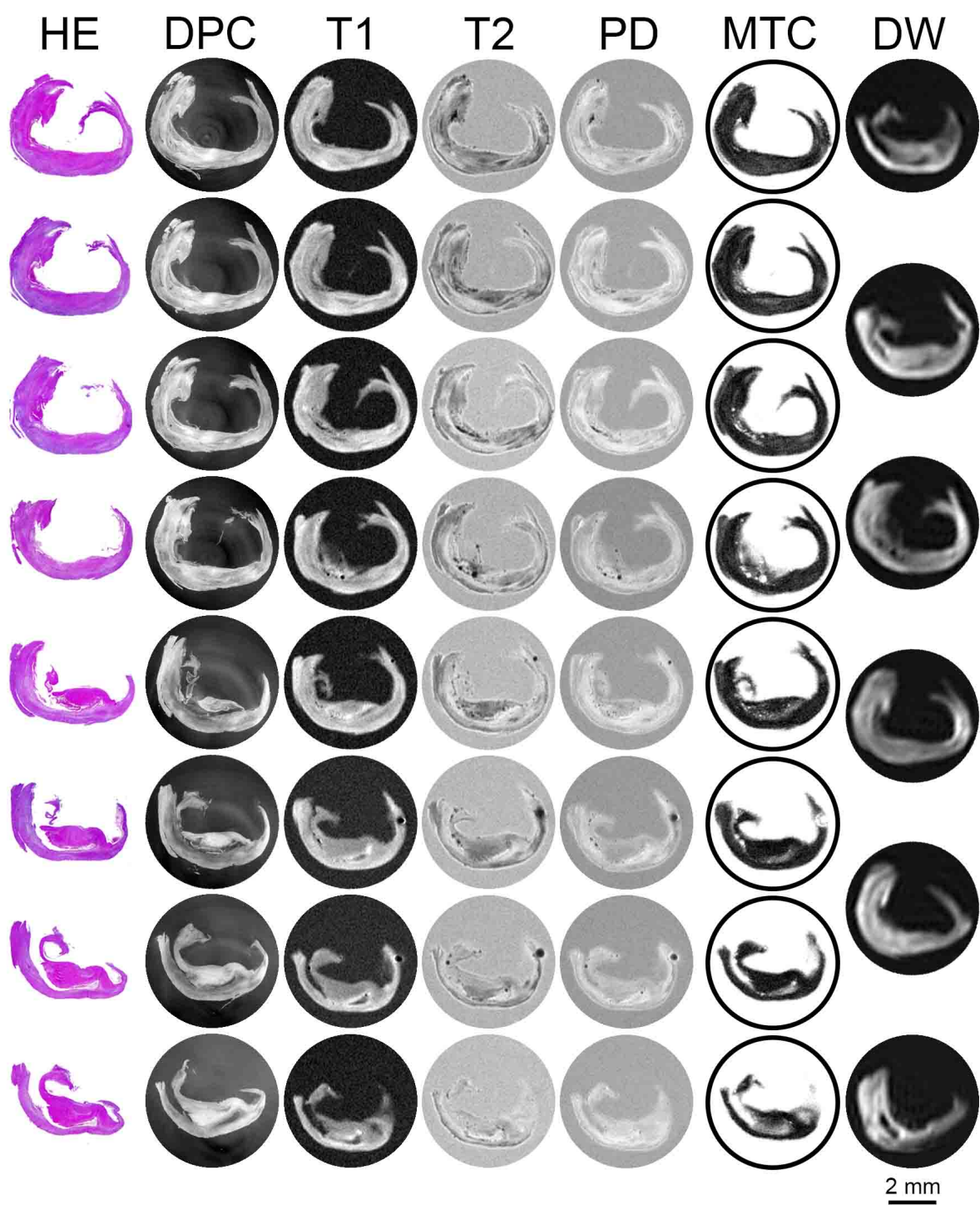


Figure S1 Serial, matched images of hematoxylin and eosin staining (HE), differential phase contrast (DPC) tomography and multicontrast MRI (T1-, T2-, proton density (PD)-weighted images, magnetization transfer contrast (MTC), diffusion-weighted (DW) images) of the stable plaque.

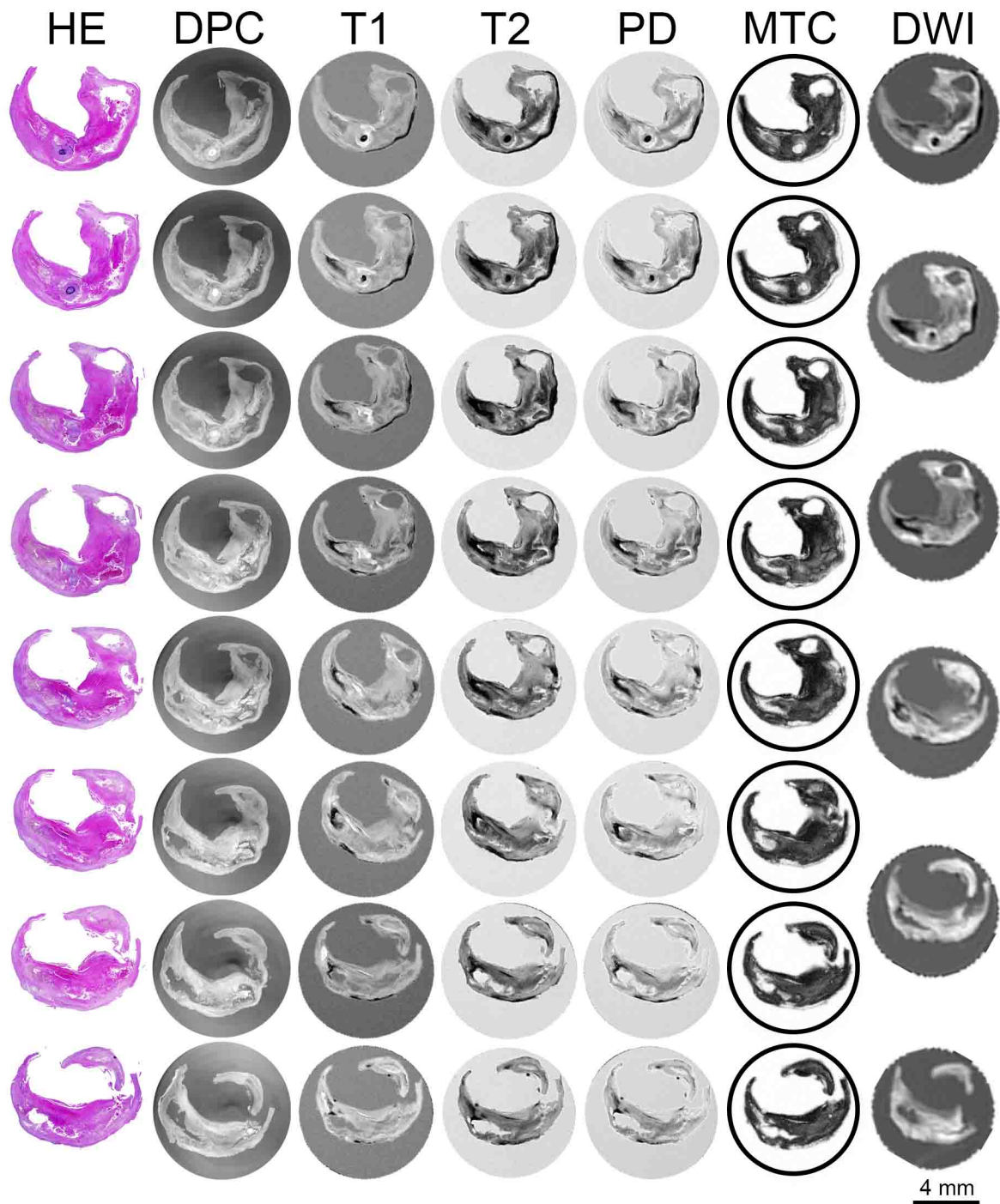


Figure S2 Serial, matched images of hematoxylin and eosin staining (HE), differential phase contrast (DPC) tomography and multicontrast MRI (T1-, T2-, proton density (PD)-weighted images, magnetization transfer contrast (MTC), diffusion-weighted (DW) images) of the unstable plaque.

<https://www.dropbox.com/s/a8h1bgobcd4dc7u/Video%20S1.gif?dl=0>

Video S1 Animated GIF file of the stable plaque after DPC tomography.

<https://www.dropbox.com/s/q4u0u0622b540kl/Video%20S2.gif?dl=0>

Video S2 Animated GIF file of the unstable plaque after DPC tomography.